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# A Consensus Linkage Map of the Chicken Genome

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A consensus linkage map has been developed in the chicken that combines all of the genotyping data from the three available chicken mapping populations. Genotyping data were contributed by the laboratories that have been using the East Lansing and Compton reference populations and from the Animal Breeding and Genetics Group of the Wageningen University using the Wageningen/Euribrid population. The resulting linkage map of the chicken genome contains 1889 loci. A framework map is presented that contains 480 loci ordered on 50 linkage groups. Framework loci are defined as loci whose order relative to one another is supported by odds greater than 3. The possible positions of the remaining 1409 loci are indicated relative to these framework loci. The total map spans 3800 cM, which is considerably larger than previous estimates for the chicken genome. Furthermore, although the physical size of the chicken genome is threefold smaller than that of mammals, its genetic map is comparable in size to that of most mammals. The map contains 350 markers within expressed sequences, 235 of which represent identified genes or sequences that have significant sequence identity to known genes. This improves the contribution of the chicken linkage map to comparative gene mapping considerably and clearly shows the conservation of large syntenic regions between the human and chicken genomes. The compact physical size of the chicken genome, combined with the large size of its genetic map and the observed degree of conserved synteny, makes the chicken a valuable model organism in the genomics as well as the postgenomics era. The linkage maps, the two-point lod scores, and additional information about the loci are available at web sites in Wageningen ([http://www.zod.wau.nl/vf/research/chicken/frame\\_chicken.html](http://www.zod.wau.nl/vf/research/chicken/frame_chicken.html)) and East Lansing (<http://poultry.mph.msu.edu/>).

The chicken is increasingly becoming of great interest as an intermediate evolutionary model organism, ideally placed between mammals and more distant vertebrates as the pufferfish and zebrafish. There are a number of different reasons for this increasing interest in the chicken genome. First, the genome size is only one-third that of mammals (Tiersch and Wachtel 1991) mainly because of its low amount of repetitive sequences and reduced intron sizes (Hughes and Hughes 1995). Furthermore, It has an interesting complex genomic structure with two chromosomal subtypes—

macrochromosomes and microchromosomes (Bloom et al. 1993)—with the microchromosomes appearing to be somewhat more gene dense than the macrochromosomes, reaching densities comparable to that of the *Fugu* genome (McQueen et al. 1998; Clark et al. 1999). Second, the level of conserved synteny between chicken and humans appears to be very high (Burt et al. 1995; Hu et al. 1995; Klein et al. 1996; Jones et al. 1997; Groenen et al. 1999; Nanda et al. 1999). Third, the chicken is being studied intensively for genes affecting polygenic traits (quantitative trait loci or QTL), which drive international efforts toward detailed physical and linkage mapping in the chicken.

Although the first genetic linkage map in chicken

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was published >60 years ago (Hutt 1936), it was not until the development of large numbers of molecular markers in the last decade that the generation of linkage maps in chicken increased. In chicken, three different linkage maps were developed using three different mapping populations. The first genetic map, based completely on DNA markers, was published by Bumstead and Palyga (1992). This map, based on the Compton (C) reference population, consisted solely of restriction fragment length polymorphism (RFLP) markers. The second genetic map to be published (Levin et al. 1993, 1994) was based on the East Lansing (EL) reference population and consisted primarily of RFLPs, random amplified polymorphic DNA (RAPD) markers, and chicken repeat element 1 (CRI) markers. Since then, both populations have been used to map a considerable number of microsatellite markers (Cheng et al. 1995; Crooijmans et al. 1997; Gibbs et al. 1997) and AFLP markers (Knorr et al. 1999) as well. The third map (Groenen et al. 1998, Herbergs et al. 1999) was based on a large F<sub>2</sub> population and consisted solely of microsatellite and amplified fragment length polymorphism (AFLP) markers. Increasing marker densities and increased initiatives in physical mapping in chicken have necessitated the need of a single consensus linkage map in chicken. Because all three maps have many markers in common, this goal has become feasible for the large and intermediate-sized chromosomes.

In this paper we describe the integration of all available data of the three mapping populations, resulting in a consensus linkage map of the chicken genome comprised of 50 linkage groups, with a total of 1889 loci.

## RESULTS AND DISCUSSION

### Linkage Maps

The genotyping data from the three chicken mapping populations were combined and analyzed simultaneously using the CRIMAP linkage program. Contributions of genotyping data were made from laboratories that have been using the EL and C reference populations and from the Animal Breeding and Genetics Group of the Wageningen University using the Wageningen/Euribrid (WAU) population. A complicating factor for the integration of all maps in chicken is the fact that not all types of markers are evenly distributed over the macro- and microchromosomes, particularly the low abundance of microsatellites on the microchromosomes (Primmer et al. 1997). Consequently, many of the small linkage groups do not have a marker in common, making the integration impossible at present. Furthermore, linkage groups C15 and C20 had only one marker in common with the corresponding linkage groups in the WAU and EL data sets, and as a consequence, the other loci from C15 and C20 could

not be positioned very precisely (Fig. 1, linkage groups E18C15W15 and E49C20W21).

The total number of different loci that have been typed on at least one of the three mapping populations was 2019. However, a relatively large proportion of these markers was either unlinked (95) or could not be positioned clearly on the linkage maps (35) and therefore were omitted from the final map shown in Figure 1. A large proportion of the omitted markers (89) are AFLP markers typed on the WAU population (Herbergs et al. 1999).

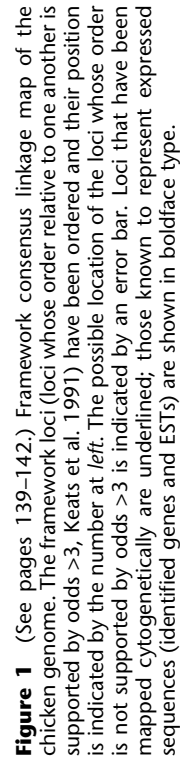
The resulting linkage map of the chicken genome contains 1889 loci (Table 1). The framework map contains 480 loci ordered on 50 linkage groups (Fig. 1). The possible positions of the remaining 1409 loci are indicated relative to these framework loci. When all linkage groups are taken into account, the total length of the linkage map is 4000 cM. However, it is expected that several of the smaller EL, C, and WAU linkage groups belong to the same chromosomes. If we correct for this fact, then the minimal length of the chicken consensus linkage map is ~3800 cM, which still is considerably larger than the previous estimates 2600–3000 cM for the chicken genome (Rodionov et al. 1992; Burt et al. 1995).

Although there are large differences in length between many of the EL (male) and the C linkage groups (female), these variations are most likely the result of differences between the lines used and typing errors in some of the RFLP markers in the C map. The differences in length between the male and female maps

**Table 1.** Number and Type of Loci on Chicken Linkage Maps

	WAU	EL	C	Consensus Map
<i>A. Loci</i>				
WAU	1011	290	119	923
EL	290	1068	195	1050
C	119	195	447	428
Type I loci	93	252	107	350
Linkage groups	34	42	36	50
<i>B. Markers</i>				
Microsatellite	573	479	190	801
Minisatellite	—	34	30	40
RFLP	—	92	191	244
AFLP	350	202	—	552
SSCP	—	50	15	59
ASO	—	71	1	71
RAPD	—	65	—	65
CR1	—	47	—	47
Classical	—	10	2	10

(A) The total number of different loci analyzed on the Wageningen (WAU), East Lansing (EL), and Compton (C) linkage maps, as well as the number of loci that are shared among the different maps. (B) The different types of markers on the different chicken linkage maps. Included are loci located on the consensus linkage map, as in Fig. 1.







**Figure 1** (Continued)



based on the WAU population generally are small with an overall difference between male and female maps of only 1.15%. Because of the much larger number of informative meioses within the WAU population, the framework of the consensus map is mainly built up by microsatellites typed on this population. Therefore, for the consensus map, the differences seen between the size of the male and female maps are similar as those described for the WAU map (Groenen et al. 1998).

### Discrepancies Between the Different Maps

Discrepancies for seven loci were observed in map locations in the original maps from the three populations used. (1) *G6PD* has been mapped on E1 and CW, and (2) *EIF4A2* has been mapped on E36 and C3. As these are both RFLPs, it is possible that different bands were scored in each population. With regard to *EIF4A2*, it is noteworthy that its location within the EL data is supported by the comparative mapping data (Fig. 2). In the EL data *EIF4A2* maps to E36, close to the *SNON* and *TFR* genes. In human, all three genes are located on the q arm of chromosome 3. However, given the discrepancies in the data, these two loci have not been included on the map. (3) *LEI0144* has been mapped on chromosome 4 (WAU) and chromosome Z (EL); (4) *MCW0066* has been mapped on chromosome 2 (C), E30 and W10; (5) *MCW0166* has been mapped on chromosome 2 (WAU) and chromosome 4 (EL). (6) The *BAT8* gene was mapped by a single laboratory (Spike and Lamont 1995) to the end of chromosome 4 (EL) and chromosome 16, the chromosome to which the major histocompatibility complex (MHC) class I and II genes have been mapped. Finally, (7) The  $\alpha$ -tubulin gene (*TUBA*) was mapped to C15 and to E22. Because this gene has been mapped by two different laboratories using different methods, it is likely that two different loci of the  $\alpha$ -tubulin gene family have been mapped. These two genes have been included in the map as *TUBAa* and *TUBAb*, respectively.

To resolve the discrepancies between the microsatellites, these markers were retyped. Typing of *LEI0144* on the EL population by M. Groenen and coworkers, (unpubl.) showed that this gene also mapped to chromosome 4, which is in agreement with the WAU data. Similarly, retyping of this marker by T. Burke and coworkers, (unpubl.) confirmed their previous results of *LEI0144* mapping to the Z chromosome. A possible explanation for this discrepancy is that the typings are done by the two groups using different primers [Crooijmans et al. (1997) and Gibbs et al. (1997), respectively]. This marker has therefore been included on both locations on the map as *LEI0144a* and *LEI0144b*. The new typing results showed clearly that the previous assignment of *MCW0066* to C chromosome 2 and of *MCW0166* to EL chromosome 4 are not correct and

that these two markers are on linkage group E30C14W10 and chromosome 2, respectively.

In addition, although there were no conflicting data between them, the mapping results of two loci for the three different populations were rather unexpected or somewhat unlikely. Marker *LEI0192* appeared to be unlinked in the EL population, whereas it is mapped to chromosome 6 using the WAU population and to linkage group C21 using the C population. Similarly, marker *HUJ0005* has been mapped to chromosome 6 on the WAU map and it is unlinked in the other two maps. Moreover, both markers are linked tightly to each other and located at the end of chromosome 6 of the WAU map, whereas they are unlinked in both of the other maps. Flipping of the typing phase of *HUJ0005* in the original MAPMANAGER file of the EL population results in linkage to several markers on chromosome 6 but not to *LEI0192*. Flipping of the phase had no effect in these markers in the C data set. Physical mapping of a BAC clone containing *LEI0192* confirmed its location at the end of chromosome 6 (V. Fillon and A. Vignal, pers. comm.).

Two additional discrepancies are observed between the consensus linkage map and the cytogenetic map. The *CYP19* gene was mapped on linkage group E029C09W09, whereas it was mapped to chromosome 1 on the cytogenetic map (Tereba et al. 1991). Comparative mapping data support the location of this gene on linkage group E29C09W09. On the linkage map the *MAX* gene maps to chromosome 4, whereas cytogenetically it was mapped to chromosome 5p (Nanda et al. 1997). In this case, the comparative mapping data support the cytogenetic location of this gene on chromosome 5, where several other human genes have been mapped that are located on chromosome 14q.

Finally, *LEI0229* was mapped to both the Z chromosome (EL) and the W chromosome (C). The most likely explanation is that *LEI0229* maps to one of the pseudoautosomal regions of the chicken Z chromosome (Fridolfsson et al. 1998).

### Anchoring Linkage Groups to Chromosomes

In addition to the integration of the three linkage maps, eventually these maps will have to be integrated with the physical map in chicken. However, the integration of the physical and genetic maps presents considerable difficulties and has proceeded at a slower pace, as a result (Morisson et al. 1998). The chicken karyotype is composed of  $2n = 78$  chromosomes which, according to their size, are classified as macro- and microchromosomes (Bloom et al. 1993). Due to the presence of microchromosomes in chicken, a standard karyotype could only be established for the eight large macrochromosomes and the two sex chromo-



Chromosome 1		
SMOH	7q31-q32	-
GAT/L3	10p15	2
NRCAM	7q31.1-31.2	12
LYZ	12	10
MGF	12q22	10
IGF1	12q22-q23	10
GNRII	8p21-p11.2	14
NAGA	22q13-qter	15
ADSL	22q13.1	15
H5	22q13.1	15
LGALS4	22	7
MGP	12p	6
LDHB	12p12.2-q12.1	6
GAPD	12p13	6
TCRB	7q35	6
HSD3B	1p13.1	3
G6PD	Xq28	-
CRYAA	21q22.3	17
ZFX/Y	Xp22.1	X
LAMP	13q34	8
P2Y5	13q14.3	-
RB1	13q14.12-q14.3	14
RAB6	2q14-21	9
PGR	11q22.1-22.3	9
FUT4	11q21	9
WNT11	11q13.5	7
HBB	11p15.4	7
NFYB	12q22-24.1	-

Chromosome 2		
ACVR2	3p22-p21.3	-
SHH	7q36	5
EN2	7q36	5
VIM	10p13	2
MRC1	10p13	2
NPY	7p15.1	6
RARB	3p24.3-p24.2	14
CP49	3q21-q25	-
TGFBR1	9q33-q34.1	4
BMP6	6pter-qter	13
BCL2	18q21.33	1
YES1	18p11.31-p11.22	5
MYL	18p11.3	5
ZFP161	18pter-p11.21	17
ADCYAP1	18p11	-
PENK	8q11.23-q12	4
CALB1	8p12-q11.2	4
CA2	8q22	3
TRHR	8q23	15
MYC	8q24.12-q24.13	15

Chromosome 4		
HMG14A	21q22.3	16
UBE2A	Xq24-q25	X
PGK1	Xq13.3	X
FMR1	Xq27.3	X
MADH1	4q28	13
GC	4q12-q13	5
ATB	4q11-q13	5
SPP1	4q11-q21	5
ANX5	4q26-q28	3
NFKB1	4q24	3
BNC1	4	5
MSX1	4p16.3-p16.1	5
TGFBR2	3p22	9
CD8A	2p12	6
TNC	9q32-q34	4

Chromosome 3		
BMP2	20p12	2
ADPRT	1q42	1
TGFB2	1q41	1
ACIN2	1q42-q43	13
ARIH2	16p13.3	5
TIMX1	4p16.1	-
MPR1	6q25.3	17
TCP1	6q25.3-q26	17
ESR	6q25.1	10
VIP	6q24-27	10
MYB	6q23.3-24	10
PLN1	6q22.1	10
FYN	6q21	10
GSTA2	6p12	9
MTF1	6q12	9
BMP5	6q12-13	9
EEF1A	6q14	4
RPL18A	19q13.2-q13.3	-
ODC1	2p25	12
MYCN	2p24.3	12

Chromosome 5		
INS	1p15.5	7
TH	1p15.5	7
CAPN1	1q13	19
RYR3	15q14	2
PTAFR	1p35-p34.3	4
HTR1D	1p36.3-p34.3	4
BRF1	14q22-24	-
TGFβ3	14q24	2
HSPCAL4	14q32.3	12
CKB	14q32.3	12
DNECL	14q32.3-qter	14
BMP4	14q22-23	14

Chromosome 6		
PSAP	10q22.1	10
PDE6C	10q24	7
ACTA2	10q22-q24	7
SCD1	10q23	19

Chromosome 7		
COL3A1	2q31-q32.3	1
FN1	2q34	1
GBX2	2q37	-
NDUSF1	2q33-34	1
EEF1B	2	1
CD28	2q33	1
VIL	2q35	1
NRAMP1	2q35	1
RPL37A	2q33-37	-
TNHB	2cen-q13	-
MC6	2q14-q21	2

Chromosome 8		
GLUL	1q25	-
PLA2G2A	1q23-qter	-
PTGS2	1q25.2-25.3	1
VTG2	9p21	4
RPL5	1p33-p32	-
GGTB1	1p33-q34	4
JAK1	1p32.3-p31.3	4
DDIT	1p31.2-31.1	3

Chromosome Z		
PRLR	5p14-p13	15
GHR	5p14-p12	15
CTSL	9p22.1-q22.2	3
PTCH	9q22.3	13
CHD1	5q15-q21	17
CHRNB3	8p11.2	8
LPL	8p22	8
ALDOB	9q22.3-q31	4
XPA	9q22.3	4
GGT2	9p21-p13	4
ACOL1	9pter-pter	4

Chromosome 16		
MHC	6p21.3	17
TAP2	6p21.3	17
BAT8	6p21.3	17
RNR1	13p12	-

E36C06W08		
TFRC SNON	3q26.2-qter 3q25-27	16 3
NCL	2q12-qter	1

E29C09W09		
B2M	15q21-q22.2	2
CYP19	15q21	9
POLG	15q26.1	-
AGC1	15q26.1	-
IGF1R	15q26.1-qter	7
RPL4	15q	-
GNRHR	4q21.2	5

E30C14W10		
CMAF2	16	8
H2AZ	6p21.3	-
CCNE	19q12-q13	7

E21E31C25W12		
MYH@	17pter-p11	11
H3F3B	17q25	11
FASN	17q25	11
RAC3	17q24-qter	11
NME2	17q21	11
IILF	17q22	11

E48C28W13W27		
MSX2	5q34-q35	13
SPARC	5q31-33	-
POU4F3	5q31	18
SPOCK	5q31	18
CDX1	5q31-33	18
CAML	5q23	13

E35C18W14		
HBA	16p13.3	11
NTN2	16p13.3	-

E18C15W15		
CRYBB1	22q11.2	5
IGLC1	22q11.2	-
MTFL2	22q11.2	10

E53C34W16		
CAMK4	5q21-q23	18
AMH	19p13.3	10

E41W17		
Ring3L	9q34	-
ABL1	9q34	2
RPL7A	9q33-q34	2
AKI1	9q34.1	2
CD39L1	9q34	2
AMBP	9q32-q33	4

E46C08W18		
HSP1	8q24.3	15
LIMK2	22q12	1

E52W19		
ACACA CRK	17q21 17p13	- 11
CASP1	11q22.2-q22.3	9

E59C35W20		
TCRA	14q11.2	14
GH	17q22-24	11
COL1A1	17q21.3-q22	11
STC4A1	17q12-q21	11

E49C20W21		
ETS1	11q23.3-q24	9
OPCML	11q23-qter	9
RPS25	11q23.3	9
POU2F3	11	9
APOA1	11q23.3	9

E16C17W22		
ARHGDTA	17q25.3	-
LUCA9	3p21.3	-
ARF2	3p21.2-21.1	-

E60E04W23		
PGA@	11q13	-
TAX1	1q32	-
TNNI2	1q32	-

E22C19W28		
GLI	12q13.2-13.3	10

E25C31		
TGFB1	19q13.2	7
RXR1	19q13.1	7

E26C13		
MCL1	1q21	3

F.32		
BMP7	20pter-qter	2
HCK	20q11-q12	2

E54		
CDC2L1	1p36	4
AGR1	1p36.3-p32	4
ENO1	1p36ter-p36.13	4
PLOD	1p36.3-p36.2	4
SLC2A1	1p35-p31.3	4

<b>E57</b>		
TP53	17p13.1	11

C24		
HSP5A	9q33-34.1	-

**Figure 2** Comparative mapping results among chicken, man, and mouse. The order of the loci is according to the linkage map shown in Fig. 1. The second column in each linkage group shows the location of the loci on the human cytogenetic map according to Genome Data Base (<http://www.gdb.org/>); the third column shows the map location in the mouse. Blocks of conserved synteny between chicken and man and between chicken and mouse are shaded.

somes (International Committee for the Standardization of the Avian Karyotype).

Many markers on the consensus map have been mapped cytogenetically as well, allowing the integration of the linkage map with the physical map. These loci are underlined in Figure 1. Because only a standard karyotype has been established for the macrochromosomes, only the linkage groups of these larger chromosomes could be assigned to their corresponding chromosomes. For historical reasons, the microchromosomes containing the MHC is named chromosome 16.

To enable identification of the microchromosomes, a set of large insert clones is being developed that can be used as tags in two-color fluorescence *in situ* hybridization (Fillon et al. 1998). Polymorphic markers have been developed for many of these large insert clones (Morisson et al. 1998; P.A. Thomson and T. Burke, unpubl.), which will allow the assignment of the linkage groups to the corresponding microchromosomes as well. Furthermore, additional large insert BAC clones have been isolated using microsatellites from the small linkage groups, which provides additional probes for the identification of the microchromosomes (R. Crooijmans, V. Fillon, M. Groenen, and A. Vignal, unpubl.).

### Comparative Gene Mapping

Genetic markers within or adjacent to known genes have been classified as type I markers (O'Brien 1991). The inclusion of type I markers on the linkage map makes it possible to access the mapping information that is available in densely mapped species such as humans and mice. Currently, the consensus map contains 350 markers within expressed sequences, 235 of which represent identified genes or sequences that have significant sequence identity to known genes. These loci are shown in boldface type in Figure 1. The orthologs of 204 of these 235 genes have also been mapped in human (Fig. 2). The comparative mapping data based on the consensus linkage map show a considerable amount of chromosomal conservation retained between man and chicken during evolution. This is in sharp contrast with the comparative mapping data between chicken and mouse, in which the amount of chromosomal conservation is considerably lower. Similar results are obtained for the comparisons between the genomes of different mammals, indicating that there have been extensive rearrangements during the evolution of the mouse genome and at a much higher rate than in birds or the other mammals (Andersson et al. 1996). Based only on the linkage data presented in this paper, at least 87 different chromosomal regions can be identified between man and chicken (Fig. 2). For many of these chromosomal regions, physically mapped genes provide additional evidence for the observed conservation of linkage (Burt et

al. 1995; Andersson et al. 1996; Nanda et al. 1999; Burt et al. 1999). Furthermore, the physically mapped genes have identified additional conserved regions between the genomes of chicken and man. Based on this number of conserved chromosomal regions, it has been calculated that the number of autosomal conserved segments shared between the chicken and human genomes is probably <100 (Burt et al. 1999). This level of conservation of synteny between chicken and human, in combination with the threefold more compact genome of the chicken, makes it an excellent evolutionary model organism in addition to Fugu, mouse, and rat. This is particularly true for the microchromosomes, which appear to be somewhat more gene dense than the macrochromosomes, thereby reaching gene densities close to that of Fugu (Angrist 1998; McQueen et al. 1998; Clarke et al. 1999). Furthermore, the higher level of conservation of genome organization (Gilley et al. 1997; Reboul et al. 1999) and the easy accessibility of the chicken as an experimental animal in studies regarding complex polygenic traits are additional features favoring the chicken over other models such as Fugu.

Although the number of loci that are available for comparative mapping are still too limited to draw detailed conclusions, it is noteworthy that several of the small linkage groups in chicken, which most likely represent different microchromosomes, seem to represent, in almost their entirety, large fragments of specific human chromosomes (e.g., E29C09W09, E21E31C25W12, E48C28W13W27, E41W17, E54, E49C20W21, and chromosome 7).

### Future Directions

The current map contains 801 microsatellite markers, which are the markers of choice for whole genome scans. However, the marker density is only sufficiently high for the macrochromosomes and a subset of the microchromosomes. Therefore, many more microsatellites are still needed to obtain (near) complete genome coverage in these kinds of studies. The integration of all the linkage maps and the cytological map in chicken is the first necessary step toward achieving this goal by identifying those regions that are particularly devoid of microsatellite markers. The major drawback, however, is the relatively low abundance of microsatellites on many of the microchromosomes (Primmer et al. 1997). Currently, increasing efforts are being put into the development of physical maps for several regions of the chicken genome, for example, Chromosome 16 (N. Bumstead, unpubl.) and linkage groups E29C09W09 and E53C34W16 (R. Crooijmans and M. Groenen, unpubl.). This has become feasible through an increased number of loci on the linkage map and because of the development of publicly available chicken YAC (Toye et al. 1997) and BAC (R.P.M.A.

Crooijmans, J. Vrebalor, R.J.M. Dijkhof, J.J. van der Poel, and M.A.M. Groenen, in prep.; J. Dodgson, unpubl.) libraries. It is to be expected that physical maps eventually will become available for all chicken chromosomes. This in turn will make the targeted development of microsatellite markers possible for those regions that currently lack any such markers, thereby allowing the characterization of these regions in QTL studies as well.

## METHODS

### Mapping Populations

The three mapping populations have been described in detail previously. Briefly, the EL population (Crittenden et al. 1993) consists of 52 BC1 animals derived from a backcross between a partially inbred jungle fowl line and a highly inbred white leghorn line. The C population (Bumstead and Palyga 1992) consists of 56 BC1 animals derived from a backcross between two inbred white leghorn lines that differed in their disease resistance. The WAU population (Groenen et al. 1998) consists of 456 F<sub>2</sub> animals from a cross between two broiler dam lines originating from the white Plymouth Rock breed.

### Markers

A detailed description of all individual loci, including their references and the number of informative meioses, is available at the web site of the Animal Breeding and Genetics Group in Wageningen ([http://www.zod.wau.nl/vf/research/chicken/frame\\_chicken.html](http://www.zod.wau.nl/vf/research/chicken/frame_chicken.html)) and East Lansing (<http://poultry.mph.msu.edu/>)

### Linkage Analysis

For each of the linkage groups, the genotyping data of the three populations were combined into a single file. To analyze the genotyping data of the backcross populations, together with data from the WAU population, the genotypes for these two populations were recoded as either being 1:1 (homozygous) or 1:2 (heterozygous). The combined data therefore consisted of 12 individual families, 1 EL, 1C, and 10 W. Linkage analysis was performed using CRIMAP version 2.4 (Green et al. 1990). Initially, a two-point linkage analysis was performed in which all markers were analyzed against each other. Tables containing all two-point lod scores for all markers are available at the web site of the Animal Breeding and Genetics Group in Wageningen. When possible, markers that had been typed on all three maps were used to start building the map using the CRIMAP-BUILD option. Finally, the order of the framework loci was checked using the CRIMAP-flips5 function.

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